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TRANS	MITTAL LETTER TO THE UNITED STATES	UNMC 63124				
DES	IGNATED/ELECTED OFFICE (DO/EO/US)	U.S. APPLICATION NO. (If known, see 37 CFR 1.5)				
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PCT/US99/07		09 April 1998 (9.04.98)				
TITLE OF INVENTION Improved Live Attenuated Viruses for Use as Vectors or Vaccines						
APPLICANT(S)	FOR DO/EO/US TRACY, Steven M. and CHAPMAN, Nora M.					
Applicant herewit	h submits to the United States Designated/Elected Office (DO/EO/US) the follow	owing items and other information:				
1. X This is	a FIRST submission of items concerning a filing under 35 U.S.C. 371.					
	a SECOND or SUBSEQUENT submission of items concerning a filing under					
examir	press request to begin national examination procedures (35 U.S.C. 371(f)) at an action until the expiration of the applicable time limit set in 35 U.S.C. 371(b) are per Demand for International Preliminary Examination was made by the 19th mc	nd PCT Articles 22 and 39(1).				
5. X A cop	y of the International Application as filed (35 U.S.C. 371(c)(2))					
a	is transmitted herewith (required only if not transmitted by the Interr	national Bureau).				
b. X	has been transmitted by the International Bureau.					
C. L	is not required, as the application was filed in the United States Rece					
	slation of the International Application into English (35 U.S.C. 371(c)(diments to the claims of the International Application under PCT Article					
a. C	are transmitted herewith (required only if not transmitted by the Inter					
b. С	have been transmitted by the International Bureau.	mational bucau).				
. c. [have not been made; however, the time limit for making such amend	ments has NOT expired.				
d. [have not been made and will not be made.	·				
8. A tran	slation of the amendments to the claims under PCT Article 19 (35 U.S.	C. 371(c)(3)).				
9. X An oa	th or declaration of the inventor(s) (35 U.S.C. 371(c)(4)).					
	slation of the annexes to the International Preliminary Examination Rep. S.C. $371(e)(5)$).	port under PCT Article 36				
Items 11, to 1	6. below concern document(s) or information included:					
	formation Disclosure Statement under 37 CFR 1.97 and 1.98.					
	signment document for recording. A separate cover sheet in compliance	with 37 CFR 3.28 and 3.31 is included.				
13. A FIR	ST preliminary amendment.					
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must also by filed (Note 37 CFR 1.9, 1.27, 1.28).			s	203.00				
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Steven M. Tracy, et al.

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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant or Patentee: Steven M. Tracy and Nora M. Chapman

Application or Patent No.: NOT YET ASSIGNED

Filed or Issued: CONCURRENTLY HEREWITH

For: Improved Live Attenuated Viruses for Use as Vectors or Vaccines

VERIFIED STATEMENT (DECLARATION) SUPPORTING ANOTHER'S CLAIM FOR SMALL ENTITY STATUS (37 CFR \$1.9(f) AND \$1,27(d)) - NONPROFIT ORGANIZATION

for purposes of paying reduced fees with rega	rd to the above-identified invention described in
(X) the specification filed herewith	
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hereby declare that I am an official empower	red to act on behalf of the nonprofit organization identified below:
FULL NAME OF ORGANIZATION:	TYPE OF ORGANIZATION
OLL RULING OF GROOM	[X] University or other institution of Higher education
BOARD OF REGENTS OF THE	[] Tax exempt under U.S. Internal Revenue Code [26 USC\$ 501(a) and
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of paying reduced fees under section 41(a) an	n identified above qualifies as a nongroff organization as defined in 37 CFR \$1.9te) for purposes of (b) of Title 35, United States code to the above-identified invention. law have been conveyed to and remain with the nonprofit organization with regard to the above-
is listed below* and the organization knows of quality as an independent inventor under 37 C	are not exclusive, each individual, concern or organization known to have rights to the invention f no rights to the invention being held by any person, other than the inventor, who could not TFR §1.8(4) if that person had made the invention, or by any concern which would not qualify as a or by a nonprofit organization under 37 CFR §1.8(a).
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may jeopardize the validity of the application, any patent issuing thereon, or any patent to which this varified statement is directed.

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IMPROVED LIVE ATTENUATED VIRUSES FOR USE AS VECTORS OR VACCINES

Pursuant to 35 U.S.C. §202(c), it is acknowledged that the U.S. Government has certain rights in the invention described herein, which was made in part with funds from the National Institutes of Health, Grant No. R21-AI42153.

This application claims priority to U.S. Provisional Application Serial No. 60/081,138, filed April 9, 1998, the entirety of which is incorporated by reference berein.

FIELD OF THE INVENTION

The present invention relates generally to the fields of molecular biology and virology. More specifically, the invention provides modified viruses for use as vaccines or vectors, which are improved in their ability to retain engineered attenuations.

BACKGROUND OF THE INVENTION

Several publications and patents are referenced in this application to describe the state of the art to which the invention pertains. Each of these publications or patents is incorporated by reference herein.

RNA viruses and retroviruses use viral encoded

25 polymerases which have a low fidelity, thereby increasing
the number of mutations that occur during replication of
the viral genome. This low fidelity results in a virus
population that contains a large number of variants. The
large variation enables these viruses to rapidly evolve

30 to adapt to a changing environment, such as a reactive
immune system, or to lose attenuating mutations
introduced to limit replication of the virus in the host.

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The RNA-dependent RNA polymerase (RDRP), has an error rate of about 10⁻⁴ (i.e., one error is introduced per every 10,000 nucleotides). Thus, each time the average RNA virus genome (ranging in size from 7,400 to ~20,000 bases) is replicated in either the positive or negative strands, at least one new random error is introduced. Reverse transcriptase (RT) has a similar error rate.

A good example of the negative impact of such a mutation rate can be observed in the Sabin poliovirus vaccine, which is a live, attenuated virus carrying a single primary attenuating mutation in the 5' nontranslated region. Within 3-5 days post-vaccination in children, the poliovirus shed in the stool has reverted to a virulent genome and the major attenuating mutation site has changed back to wild-type.

The main factor contributing to high mutation rates in these viruses is the absence or low efficiency of proofreading or repair activities associated with RDRPs. Structurally, the RDRPs are similar to one another (for instance, HIV reverse transcriptase and poliovirus RNA polymerase are quite similar). The poliovirus RNA polymerase is shaped like a hand that is making an effort to hold a cup or glass, fingers somewhat together and curled, thumbs apart, with a palm in between (see, e.g., Hansen, J.L., A.M. Long and S.C. Schulz (1997); Structure $\underline{5}$: 1109-1122). The palm region contains the active site for both the poliovirus and HIV polymerases, and the region contains similar motifs in both enzymes. In HIV strains from patients who no longer respond to nucleotide therapy, mutants have been isolated that (a) have mutations in the palm region, (b) show many-fold (e.g., 3-49 fold) better fidelity, and (c) replicate the genome at a reduced rate.

35 Clearly, the aforementioned high reversion rate

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in RNA viruses and other viruses utilizing RDRPs or RTs, which is caused by their low-fidelity polymerases, is detrimental to their utility as vaccines or as vectors for delivery of other genes of interest. Thus, the field of viral vaccines and vectors would be vastly improved through the development of improved vectors encoding _ RDRPs or RTs with greater replicative fidelity, such that attenuations introduced into the viral genomes are retained for longer periods of time, without reversion to wild-type virus.

SUMMARY OF THE INVENTION

The present invention provides novel modified viral genomes that encode RDRPs and RTs whose activity is altered, e.g., by having improved fidelity as compared with their unmodified counterparts. When combined with additional attenuation mutations, these genomes and their encoded viruses are superior to those currently available for use as vaccines or as vectors for delivery of other genes of therapeutic or diagnostic value.

In one aspect of the invention, there is provided a virus genome that encodes an RNA-dependent polymerase, the genome being modified to produce an attenuated virus, the genome further comprising at least one pol gene modification, which results in a decreased reversion rate from attenuated virus to non-attenuated virus as compared with an equivalent virus genome without the pol gene modification. The RNA-dependent polymerase can be an RNA polymerase or a reverse transcriptase. Preferably it is an enterovirus genome, more preferably a coxsackievirus genome. The decreased reversion rate can be the result of a variety of alterations in the polymerase, such as a decrease in rate of polymerase activity, which itself may be caused by the enzyme having increased fidelity as compared with a polymerase from a

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virus genome that does not comprise the pol gene modification.

According to another aspect of the invention, a viral vector for delivering a heterologous nucleic acid to a target cell, tissue or organ is provided. The vector comprises the aforementioned modified virus genome, and further comprises at least one cloning site for insertion of an expressible heterologous nucleic acid, such as an antigen or a biologically active molecule.

According to another aspect of the invention, the modified virus may be used as a live, attenuated vaccine for prevention of infection by that virus.

In a preferred embodiment of the present 15 invention, the polymerase-modified viral genome described above is a coxsackievirus genome, preferably a coxsackievirus 3B (CVB3) genome, modified to produce an attenuated virus. In the CVB3 genome, the pol gene modification preferably comprises a mutation at a 20 position on the genome encoding glycine 328. The attenuation mutation of the coxsackievirus genome preferably is in a transcription regulatory region, such as the 5' non-translated region of the genome. The CVB3 genome of claim 16, wherein the modification to produce an attenuated virus comprises altering a transcription 25 regulatory region of the genome. Most preferably, the genome is modified by changing U to C or G, or C to G, at nucleotide position 234 of the genome to achieve the attenuation. In alternative embodiments the genome contains other attenuating mutations, or a combination of attenuating mutations, along with the pol gene modification.

Other features and advantages of the present invention will be better understood by reference to the drawings, detailed descriptions and examples that follow.

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DETAILED DESCRIPTION OF THE INVENTION

I. Definitions

Certain aspects of the present invention employ conventional molecular biology, microbiology, and recombinant DNA techniques that are well known in the art. See, e.g., Sambrook et al., "Molecular Cloning: A-Laboratory Manual (1989); "DNA Cloning: A Practical Approach," Volumes I and II (D.N. Glover ed. 1985); "Oligonuclectide Synthesis" (M.J. Gait ed. 1984); "Nucleic Acid Hybridization" [B.D. Hames & S.J. Higgins eds. (1985)]; "Transcription and Translation" [B.D. Hames & S.J. Higgins eds. (1984)]; "Animal Cell Culture" [R.I. Freshney, ed. (1986)]; "Immobilized Cells And Enzymes" [IRL Press, (1986)]; B. Perbal, "A Practical Guide To Molecular Cloning" (1984); or "Current Protocols in Molecular Biology", eds. Frederick M. Ausubel et al., John Wiley & Sons. 1999.

Therefore, if appearing herein, the following terms have the definitions set out below.

A "coding sequence" or "coding region" refers to a nucleic acid molecule having sequence information necessary to produce a gene product, when the sequence is expressed.

The term "operably linked" or "operably
inserted" means that the regulatory sequences necessary
for expression of the coding sequence are placed in a
nucleic acid molecule in the appropriate positions
relative to the coding sequence so as to enable
expression of the coding sequence. This same definition
is sometimes applied to the arrangement other
transcription control elements (e.g. enhancers) in an
expression vector.

Transcriptional and translational control sequences are DNA regulatory sequences, such as

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promoters, enhancers, polyadenylation signals, terminators, and the like, that provide for the expression of a coding sequence in a host cell.

The terms "promoter", "promoter region" or "promoter sequence" refer generally to transcriptional 5 regulatory regions of a gene, which may be found at the 5' or 3' side of the coding region, or within the coding region, or within introns. Typically, a promoter is a DNA regulatory region capable of binding RNA polymerase in a cell and initiating transcription of a downstream (3' direction) coding sequence. The typical 5' promoter sequence is bounded at its 3' terminus by the transcription initiation site and extends upstream (5' direction) to include the minimum number of bases or 15 elements necessary to initiate transcription at levels detectable above background. Within the promoter sequence is a transcription initiation site (conveniently defined by mapping with nuclease S1), as well as protein binding domains (consensus sequences) responsible for the binding of RNA polymerase.

A "vector" is a replicon, such as plasmid, phage, cosmid, or virus to which another nucleic acid segment may be operably inserted so as to bring about the replication or expression of the segment.

An "origin of replication" refers to those DNA sequences that participate in the in the initiation of DNA synthesis.

A cell has been "transformed" or "transfected" by exogenous or heterologous DNA when such DNA has been introduced inside the cell. The transforming DNA may or may not be integrated (covalently linked) into the genome of the cell. In prokaryotes, yeast, and mammalian cells for example, the transforming DNA may be maintained on an episomal element such as a plasmid. With respect to eukaryotic cells, a stably transformed cell is one in

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which the transforming DNA has become integrated into a chromosome so that it is inherited by daughter cells through chromosome replication. This stability is demonstrated by the ability of the eukaryotic cell to establish cell lines or clones comprised of a population of daughter cells containing the transforming DNA. A "clone" is a population of cells derived from a single cell or common ancestor by mitosis. A "cell line" is a clone of a primary cell that is capable of stable growth in vitro for many generations.

A "heterologous" region of a nucleic acid construct is an identifiable segment of the nucleic acid molecule within a larger molecule that is not found in association with the larger molecule in nature. Thus, when the heterologous region encodes a mammalian gene, the gene will usually be flanked by DNA that does not flank the mammalian genomic DNA in the genome of the source organism. In another example, coding sequence is a construct where the coding sequence itself is not found in nature (e.g., a cDNA where the genomic coding sequence contains introns, or synthetic sequences having codons different than the native gene). Allelic variations or naturally-occurring mutational events do not give rise to a heterologous region of DNA as defined herein.

The terms set forth below, relating to the biological molecules and methods of the present invention, are used throughout the specifications and claims.

The term "Coxsackie B3 virus;" or "CVB3" refers

to a specific serotype of the human coxsackie B
enterovirus of the family *Picornaviridae*, genus

Enterovirus. The CVB3 genome is characterized by a single
molecule of positive sense RNA which encodes a 2,185
amino acid polyprotein.

35 The term "attenuated" refers to a virus that is

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modified to be less virulent (disease-causing) than wildtype virus.

The term "basic CVB3/0 genome" refers to the modified Coxsackievirus B3 as reported by Chapman, N.M., et al, Arch. Virol. 122:399-409 (1994).

The term "viral protease" or "viral encoded protease" refers to viral encoded enzymes that degrade proteins by hydrolyzing peptide bonds between amino residues. Some such proteases recognize and cleave at only specific sequences.

The term "open reading frame" refers to a length of RNA or DNA sequence, between a translation start signal (e.g., AUG or ATG) and any one or more of the known termination codons, which encodes one or more polypeptide sequences.

The term "RNA-dependent polymerase (RDP) refers to a viral polymerase that transcribes either RNA or DNA from an RNA template. The RNA viruses described herein produce an RNA-dependent RNA polymerase (RDRP). The retroviruses described herein produce an RNA-dependent DNA polymerase, often referred to as a "reverse transcriptase (RT)"

The term "viral vector" refers to a virus that is able to transmit foreign or heterologous genetic

25 information to a host. This foreign genetic information may be translated into a protein product, but this is not a necessary requirement for the foreign information.

The term "capsid coding region" refers to that region of a viral genome that contains the DNA or RNA code for protein subunits that are packaged into the protein coat of the virus particle.

II. Compositions and Methods

RNA viruses and retroviruses use viral encoded polymerases RNA-dependent polymerases (RDPs) which have a

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low fidelity, thereby increasing the number of mutations that occur during replication of the viral genome. The low fidelity of these RDPs negatively impacts the effectiveness of these viruses as vectors or vaccines, because they quickly lose attenuating mutations introduced to limit replication of the virus in the host. The present invention is directed to modifications of the genomes of these viruses to improve the fidelity of their RDPs, thereby reducing the rate of reversion of attenuated strains and slowing the replication rate of attenuated viruses being used as vaccines or vectors.

Any virus having utility as a vaccine or vector, which replicates by means of a RNA-dependent polymerase, is suitable for modification in accordance with the present invention. Examples of suitable viruses include, but are not limited to, enteroviruses such as coxsackieviruses, echoviruses, polioviruses and numbered enteroviruses, other RNA viruses such as flaviviruses and togaviruses, and retroviruses such as human

immunodeficiency virus (HIV), human T-cell leukemia virus (HTLV), avian sarcoma leukosis virus (ASLV), feline leukemia virus (FeLV), bovine immunodeficiency virus (BIV) and equine infectious anemia virus (EIAV), among others. Other suitable viruses can be selected from the families of mammalian viruses known to replicate using an RNA-dependent polymerase. These include picornaviruses, calciviruses, coronaviruses, retroviruses, flaviviruses,

The RDPs of the above-listed viruses are

structurally similar to one another, as demonstrated by
reference to the following representative published
literature: Jacobo-Molina A, Ding J, Nanni RG, Clark AD
Jr, Lu X, Tantillo C, Williams RL, Kamer G, Ferris AL,
Clark P, et al., (1993) Crystal structure of human
immunodeficiency virus type 1 reverse transcriptase

togaviruses, filoviruses and bunyaviruses.

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complexed with double-stranded DNA at 3.0 A resolution shows bent DNA. Proc Natl Acad Sci U S A 90(13):6320-4; Hansen JL, Long AM, Schultz SC, (1997) Structure of the RNA-dependent RNA polymerase of poliovirus. Structure 5(8):1109-22; Sousa R, Chung YJ, Rose JP, Wang BC, (1993) Crystal structure of bacteriophage T7 RNA polymerase at _ 3.3 A resolution. Nature Aug 12;364(6438):593-9; and O'Reilly EK, Kao CC, (1998) Analysis of RNA-dependent RNA polymerase structure and function as guided by known polymerase structures and computer predictions of secondary structure. Virology 252(2):287-303.

Attenuated coxsackieviruses are contemplated as being particularly useful for practice of the present invention. Methods for making modified coxsackievirus genomes with primary attenuating mutations are described in detail in co-pending U.S. Serial No. 08/812,121 and co-pending PCT Serial No. PCT US98/04291, both to Tracy and Chapman, the disclosures of which are incorporated by reference herein. Those patent documents describe a viral vector which comprises a coxsackievirus genome modified to encode an attenuated coxsackievirus, preferably a coxsackievirus B, most preferably a coxsackievirus B3

Tracy and Chapman teach that attenuation of the 25 coxsackievirus can be achieved by altering a transcription regulatory region of the genome. Preferably, the transcription regulatory region comprises a 5' untranslated region of the genome. In one instance, the 5' untranslated region is replaced with a 5' untranslated region of a non-enterovirus genome selected 30 from the group consisting of poliovirus and echovirus. In another instance, a coxsackievirus B3 genome is modified by substituting a C or G for a U at nucleotide position 234 of the genome. Another modification includes point mutations at positions nt232 and nt236, or 35 deletion entirely of nt 232-236.

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Tracy and Chapman further teach a cloning site in the coxsackievirus vector, which can be positioned between a coding sequence for the capsid protein and a coding sequence for the viral protease. Alternatively, the cloning site is positioned at the start of the genome's open reading frame, and is constructed such that the inserted expressible heterologous DNA comprises a translation start codon and a 3' sequence recognized by a viral protease.

Though coxsackievirus B3 is exemplified herein, any coxsackievirus genome is believed to be suitable for use in the present invention. This is due to the high level of structural similarity among RDPs, as discussed above, as well as substantial organizational similarity among the coxsackieviruses, and indeed among enteroviruses in general (see, e.g., Romero et al., Current Topics in Microbiology and Immunology 223: 97-152, 1997; Chapman et al., Current Topics in Microbiology and Immunology 223: 227-258, 1997; and Tracy et al., Trends in Microbiology 4: 175-179, 1996).

Attenuated strains of viruses other than coxsackieviruses are also contemplated for use in the present invention. Some of these are based on information obtained through characterization of attenuating mutations of policyirus (Minor PD, Macadam AJ, Stone DM, Almond JW, (1993) Genetic basis of attenuation of the Sabin oral policyirus vaccines. Biologicals Dec;21(4):357-63). The best defined sites of attenuation in the Sabin strains of policyirus are those in the Sabin policyirus 3 strain: the site in the 5' nontranslated region causes a reduction in initiation of translation and the sites in the capsid protein encoding region cause a change in stability of the capsid (Macadam AJ, Ferguson G, Arnold C, Minor PD, (1991) An assembly defect as a result of an attenuating mutation in

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the capsid proteins of the poliovirus type 3 vaccine strain. J Virol 65(10):5225-31). In other strains there are sites elsewhere that contribute to attenuation but all the strains contain the 5' non-translated region sites. The majority of the attenuation of all poliovirus strains is due to similar 5' non-translated region sites and capsid sites. As discussed above, attenuation of CVB3 has been made by substituting the 5' non-translated region of poliovirus for the naturally occurring coxsackievirus 5' region. Similar sorts of mutations as found in the polioviruses can be made in the coxsackievirus genome with an expectation of similar results. Alternatively, the 5' non-translated region of attenuated polio-like viruses could be substituted for the coxsackievirus 5' non-translated regions. polio-like viruses have a 5' non-translated region sufficiently similar to polioviruses (coxsackieviruses A21 and A24) that similar attenuating mutations can be made with assurance that similar attenuation will occur.

Other candidate viruses particularly suitable for attenuation in a manner similar to that shown for poliovirus and coxsackievirus, then modified in RDP function in accordance with the present invention, include other human enteroviruses such as echoviruses, animal enteroviruses, such as bovine enterovirus, as well as members of the flavivirus and togavirus families. Retroviruses that may be used are animal retroviruses capable of replicating in human cells, or animal retroviruses for use as animal vectors, as well as highly genetically engineered strains of HIV. However, as mentioned above, any virus having an RNA-dependent polymerase is considered suitable for use in the present invention.

Modified viral genomes having a reduce rate of reversion from attenuated mutant to wild-type, via

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improved transcriptional fidelity, can be selected or identified by two general approaches. In one approach, new viral mutants can be made using methods designed to force preferential mutations in the pol gene. For instance, virus is grown in the presence of ribonucleoside analogs (antiviral drugs) which inhibit _ the RDPs. Mutations (occurring naturally due to the high error rate and likely to be in the RDP-encoding region in sites similar to the drug resistance sites in HIV) which confer resistance to the ribonucleoside analogs would be selected for in this environment. Viruses with these mutations would out-compete the non-resistant viruses in the presence of the antiviral drug. Viruses are plaque-isolated after several passages in the selective media.

RDPs from mutant viruses are then obtained by RT-PCR, and those genes are used to replace the wild-type RDP coding region in a clone of the viral genome. The clones are used to raise a population of virus in an appropriate cultured cell line (e.g., HeLa cells) and the fidelity is assessed by (1) transcriptional assays that determine the frequency of incorrect rNTP incorporation, and (2) the time required for another mutation (located elsewhere in the genome, such as those we have characterized that slow growth within the viral 5' non-translated region) to revert in the virus with the mutated RDP, as compared to the wild-type RDRP virus containing the same mutation.

In another approach, specific mutations are introduced, or mutations are randomly introduced in a selected region of the RDP coding region known to influence transcriptional fidelity. For instance, mutations may be introduced in the codon at nucleotides 6893-6895 (GGT) which encodes amino acid 328 of the 3D RNA-dependent RNA polymerase, glycine. The preferred mutations are GGT -> AGT (to serine), -> TGT (to

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cysteine), -> GCT (to alanine). These mutations have been demonstrated to be viable in poliovirus but not lethal, yet with diminished polymerase activity, a phenotype that is consistent with a polymerase with greater fidelity (Jablonski SA, Luo M, Morrow CD, J Virol 65:4565-72, 1991. Accordingly, these mutations can be introduced into coxsackievirus or the other viruses listed above, with the anticipated results being similar to those observed in poliovirus.

In other embodiments, additional mutations suggested by reverse transcriptase higher fidelity mutants are mutations of the aforementioned codon to CTT (leucine) and to ATT (isoleucine) and mutations of the preceding codon (n6890-6892 TAT encoding tyrosine) to TTT (phenylalanine).

As another example, in HIV-1 reverse transcriptase, a met-val substitution at codon 184 is known to improve the fidelity of the transcriptase (Wainberg, M.A. (1997); Leukemia, Apr. 11 Supp. 3: 85-88; Oude Essink, B.B., N.K.T. Back and B. Berkhout (1997); Nucl. Acids Res. $\underline{25}$: 3212-3217). Sites in the finger region of the polymerase also have been shown to affect fidelity; these include leu-val and glu-gly substitutions at codons 74 and 89, respectively (Rubinek, T., M. Bakhanashvili, R. Taube, O. Avidan and A. Hizi (1997); Eur. J. Biochem. 247: 238-247). Similar mutations introduced into enterovirus RNA polymerases are expected to have a similar effect, due to the close structural and sequence homology among the RDPs. For instance, a site in the coxsackievirus B3 genome that corresponds to the leu-val substitution in the RT is position 194 of the CVB3 genome, nucleotides 6491-6493; and a site in CVB3 corresponding to the glu-gly substitution is position 213, nucleotides 6548-6550. In alternative embodiments, these sites are mutated to produce a modified virus with

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a reduced rate of reversion from attenuated to wild type form. $\ensuremath{\text{\text{c}}}$

Mutations are obtained by PCR, discarding nonsense mutations identified by sequence analysis. Specific mutant sequences are then used as described above to replace wild-type sequences in clones of the genomes. Virus is obtained by transfection of the genomes in a cultured cell line, and transcriptional fidelity is assayed as described above.

Consistent with the approaches described above (particularly as demonstrated by the above-mentioned assay for reversion of a second site on the genome), incorporation of sequences encoding higher fidelity RDPs will result in a virus that does not as rapidly correct mutations by the mechanism of stochastic mutation and selection of a more fit (reverted) virus population. If such mutations are primary attenuating mutations, then the resultant attenuated viral genome will be less prone to reversion to wild-type (presumably virulent). A more stable viral strain genome, for use as either a vaccine or a vector, is superior by virtue of reduced risk of reversion and resultant vaccine-related disease caused by the reverted strain. Additionally, any vaccine that is shed in feces, blood or aerosol would more likely be the attenuated strain rather than a reverted, possibly virulent, strain.

Methods for measuring the reversion rate of an attenuated virus in cultured cells and in vivo are described in Example 2. Essentially, these methods

measure delays in reversion of an attenuating mutation in viruses containing a selected RDP mutation, as compared with viruses not containing the mutation. An RDP mutant virus exhibiting any observed delay in such reversion, no matter how small, is considered to be within the scope of the present invention. Preferred embodiments of the

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invention, however, include viruses exhibiting at least a two-fold delay in reversion time, as compared to their RDP-unmodified counterparts. In particularly preferred embodiments, the delay in reversion rate would be greater; e.g., 3-5, 6-8, 9 or 10-fold or more.

The following examples are provided to describe the invention in greater detail. They are intended to illustrate, not to limit, the invention.

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EXAMPLE 1

Characterization of a CVB3 Reporter Mutant to Test for Reversion of Mutant Sites

This example describes a CVB3 reporter mutant, useful to test modifications of the *pol* gene for their effect on the rate of reversion of attenuated mutants to wild-type virus.

An infectious, but non-cardiovirulent strain of CVB3 is CVB3/0 (Chapman et al., (1994) Arch. Virol. 135(1-2): 115-130). CVB3/0 differs from the infectious, cardiovirulent strain CVB3/20 by eight amino acid sequence differences; however, the cardiovirulent phenotype of CVB3/20 is determined at a single site in the genomic 5' non-translated region (Tu et al., (1995) J. Virol. 69(8): 4607-4618).

CVB3/20 also contains an attenuating mutation at nt234 (U-G). The reversion rate of this mutation to the wild-type has been characterized *in vitro* in CVB3/20. It has been found that 234G mutation rapidly reverts to U in passage in HeLa cells. At 37°C, the reversion occurs within 3-4 passages. However, transfection of the plasmid construct at 33.5°C maintained the mutation intact, judging by sequence analysis of the 234G virus population RNA using RT/PCR and sequencing of the

amplimer obtained.

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The CVB3/20 mutant with 234G has been demonstrated to be significantly attenuated in terms of replication in both HeLa cells as well as in murine fetal heart fibroblast cultures; Because the virus is attenuated for replication in HeLa relative to the parental CVB3/20, it is expected that the 234G virus is also attenuated for replication in other human cell cultures that may be used.

The biology of mutations in the CVB3 genome in the 5mer surrounding the 234 site (nucleotides 232-236, 5'-CGUUA where 234U [the 5' U] is underlined) have been further elucidated. The inventors have observed that transfection of viruses with mutations in this 5mer at 37°C results in slow growing, nearly undetectable viruses for the first 2-3 passages in HeLa cells, then a sudden increase in titerable virus. This increase correlates directly with a reversion from the mutation to the wildtype sequence. However, if transfections are performed at 33.5°C and the stocks are passaged at 33.5°C, the virus population achieves a titerable level within passage 2-3 and also maintains the mutation judging by sequence analysis of the amplimer obtained using total viral RNA as the template. Thus, it would be expected that pol mutations introduced into the viral genome will be stable or more stable when transfected at 33.5°C, and that a delay in reversion of the 234G to U would be observed, compared to the 234G mutant alone (no pol mutation in the same virus) when the virus stocks are shifted to be passaged at 37°C.

Even though the biology of the nt234 mutation has been elucidated in CVB3/20, CVB3/0 is preferred over CVB3/20 as it confers some advantages in the manipulation of the genome but also because it plaques on HeLa cell monolayers, whereas CVB3/20 does not. The genome of CVB3/0 has a C at nt234, which confers attenuation on this clone (Tu et al., 1995, supra). However, from

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research on CVB3/20, it is known that the G mutation at nt234 confers a greater attenuation than does the C at that position. For this reason, a C-G mutation at nt234 has been introduced into the pCVB3/0 virus genome clone. The CVB3/0 nt234G attenuated mutant is particularly suited for use as a reporter for study of pol mutation _ because it is highly attenuated in terms of viral replication and the virus can be plaqued.

EXAMPLE 2 Testing CVB3 Vectors Having Modifications Expected to Produce High Fidelity Polymerases

This example describes how a CVB3 attenuated, pol modified virus, such as the one described in Example 1, is tested for rate of reversion to wild-type virus in cultured cells and in vivo. In cultured cells, the following assays are used: (a) rapidity of the appearance of cytopathic effects, using light microscopy examination of infected cultures; (b) titer of virus in infected cultures; and (c) RT-PCR amplification of the region surrounding nucleotide 234 and sequence analysis across the mutation. Concurrently, site in the pol gene which the mutation was induced is also sequenced.

Virus is passaged on 100,000 HeLa cells in a monolayer at an MOI (multiplicity of infection) of infectious virus particles (defined in TCID50) of 1. Virus is incubated with cells for 1 hour either at 37°C or 33.5°C, the virus inoculum washed off in three washes with medium, and the cells re-fed and incubated at the respective temperature. Virus can be harvested in two approaches: either at 24 hours post-inoculation or at such time that cytopathic effects (rounding, detached and floating cells, destruction of the cell monolayer) are extreme. Virus titer is ascertained at the temperature at which the stocks were propagated. As the same number

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of cells and the same volume of medium is always employed, titers from successive passages can be directly compared in terms of infectious particles per mL or as a function of the number of cells.

Virus is used as a source of viral RNA. The viral RNA is reverse-transcribed, then the cDNA amplified with primers that flank the 234 site. The amplified fragment is directly sequenced, and the identity of the nucleotide at position 234 is determined. Similarly, the region surrounding the mutation in the polymerase is amplified, and the identity of the nucleotide at the mutated site(s) is determined.

As described in Example 1, we have demonstrated that mutations in the 5mer region nucleotide 232-236 (5'CGUUA) of CVB3/20 are stable when the mutated viruses are propagated at 33.5°C but revert within 3-5 passages in cells at 37°C. At 37°C, there is a sudden 4-5 log increase in virus titer around pass 3-4 from a previously extremely low (less to much less than 500-750 TCID50/mL) titer. This sudden increase in titer correlates with the reversion of the mutated site(s) in the 5mer. No such sudden increase in titer is observed at 33.5°C and no reversion from the mutated site is observed by sequence analysis out to pass 6. These differences in reversion at the different incubation temperatures can be used to define and compare respective reversion rates in pol-mutated and non-mutated virus.

Reversion rates of attenuated viruses can also be measured in vivo. Standard tests for reversion of attenuated viral vaccines involve measuring the reversion of the attenuating mutation in excreted virus particles. Such methods have been used to measure the reversion rate of attenuated poliovirus vaccines, wherein it was discovered that attenuated sites would mutate to produce reverted virus within 24 to 48 hours in fecal samples (Dunn et al., (1985) Nature 314(6011): 548-550; Macadam

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et al., (1989) Virology 172(2): 408-414).

Measuring the reversion rate of pol mutants is first established in a suitable animal model, and thereafter can be used in humans. The mutant reporter strain is one selected to have a particularly high rate of mutation to quickly produce virus in feces that have_a reverted genome. Pol mutations introduced into these test viruses can then be tested for any delay in reversion rate.

The extent of reversion in fecal samples is measured according to standard methods. Essentially, this involves plaquing virus from feces (to obtain clonal populations derived from one virus particle), then determining the sequence at attenuating sites for each plaqued stock. Results are quantified as reverted isolates per total clonal stocks plaqued.

The present invention is not limited to the embodiments described above, but is capable of modification within the scope of the appended claims.

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We claim:

- 1. A virus genome that encodes an RNAdependent polymerase, the genome being modified to

 5 produce an attenuated virus, the genome further
 comprising at least one pol gene modification, which
 results in a decreased reversion rate from attenuated
 virus to non-attenuated virus as compared with an
 equivalent virus genome without the pol gene

 10 modification.
 - 2. The virus genome of claim 1, wherein the RNA-dependent polymerase is an RNA polymerase.
 - 3. The virus genome of claim 2, wherein the virus genome is an enterovirus genome.
 - 4. The virus genome of claim 3, wherein the enterovirus is selected from the group consisting of coxsackievirus, poliovirus polio-like virus and echovirus.
 - 5. The virus genome of claim 4, wherein the virus is coxsackievirus B, serotype 3 (CVB3).
 - 6. The virus genome of claim 1, wherein the RNA-dependent polymerase is a DNA polymerase.
- The virus genome of claim 6, from a virus
 selected from the group consisting of HIV, HTLV, ASLV,
 FeLV, BIV, and EIAV.
- The virus genome of claim 1, wherein the decreased reversion rate is caused by a decrease in rate of polymerase activity.

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- 9. The virus genome of claim 1, wherein pol gene modification results in a polymerase having increased fidelity as compared with a polymerase from a virus genome that does not comprise the pol gene modification.
- 10. The virus genome of claim 1, wherein the pol gene modification comprises a mutation resulting in an alteration of the RNA polymerase active site.
- 11. The virus genome of claim 1, having a reversion rate at least two-fold decreased as compared with an equivalent virus without the pol gene modification
- 12. A viral vector for delivering a heterologous nucleic acid to a target cell, tissue or organ, comprising the virus genome of claim 1, said genome further comprising at least one cloning site for insertion of an expressible heterologous nucleic acid.
 - 13. The vector of claim 12, comprising an expressible heterologous nucleic acid encoding an antigenic molecule.
 - 14. The vector of claim 12, comprising an expressible heterologous nucleic acid encoding a biologically active molecule.
- 30 15. A live, attenuated viral vaccine comprising the virus genome of claim 1.
- 16. A coxsackievirus 3B (CVB3) genome, modified to produce an attenuated virus, the genome
 35 further comprising at least one *pol* gene modification, which results in a decreased reversion rate from the

attenuated virus to non-attenuated virus as compared with an equivalent virus genome without the pol gene modification

- 5 17. The CVB3 genome of claim 16, wherein the decreased reversion rate is caused by a decrease in rate of polymerase activity.
- 18. The CVB3 genome of claim 16, wherein pol gene modification results in a polymerase having increased fidelity as compared with a polymerase from a CVB3 genome that does not comprise the pol gene modification.
- 15 19. The CVB3 genome of claim 16, wherein the pol gene modification comprises a mutation resulting in an alteration of the RNA polymerase active site.
- 20. The CVB3 genome of claim 19, wherein the 20 pol gene modification comprises a mutation at a position on the genome encoding glycine 328.
- 21. The CVB3 genome of claim 20, wherein the mutation results in a change in glycine 328 to cysteine 25 or alanine.
 - 22. The CVB3 genome of claim 16, wherein the modification to produce an attenuated virus comprises altering a transcription regulatory region of the genome.
 - 23. The CVB3 genome of claim 22, having its 5' untranslated region replaced with a 5' untranslated region from a heterologous genome.

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- 24. The CVB3 genome of claim 22, wherein the genome is modified by changing U to C or G, or C to G, at nucleotide position 234 of the genome.
- 5 25. A viral vector for delivering a heterologous nucleic acid to a target cell, tissue or organ, comprising the virus genome of claim 16, said genome further comprising at least one cloning site for insertion of an expressible heterologous nucleic acid.
 - 26. The vector of claim 25, comprising an expressible heterologous nucleic acid encoding an antiquenic molecule.
- 15 27. The vector of claim 25, comprising an expressible heterologous nucleic acid encoding a biologically active molecule.
- 28. The vector of claim 25, wherein the
 cloning site is positioned between a coding sequence for
 a capsid protein and a coding sequence for viral
 protease.
- 29. The vector of claim 25, wherein the
 25 cloning site is positioned at the start of the genome's open reading frame, such that the inserted expressible heterologous DNA comprises a translation start codon and a 3' sequence recognized by a viral protease.
- 30 30. A live, attenuated coxsackievirus vaccine comprising the CVB3 genome of claim 16.

I acknowledge the duty to disclose information, which is material to patentability as defined in 37, Code of Federal Regulations, Section 1.56.

CLAIM FOR BENEFIT OF PRIOR U.S. PROVISIONAL APPLICATION(S)

(35 U.S.C. Section 119(e))

I hereby claim the benefit under Title 35, United States Code, Section 119(e) of any United States provisional application(s) listed below:

PROVISIONAL APPLICATION NUMBER

FILING DATE

60/081,138

April 9, 1998

POWER OF ATTORNEY

I hereby appoint the following practitioner(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith.

APPOINTED PRACTITIONER(S)	REGISTRATION NUMBER(S)
Janet E. Reed	36,252
Alfred W. Zaher	42,248
Bruce D. George	43,631

COMBINED DECLARATION AND POWER OF ATTORNEY

(ORIGINAL, DESIGN, NATIONAL STAGE OF PCT, SUPPLEMENTAL, DIVISIONAL, CONTINUATION, OR C-I-P)

As a below named inventor, I hereby declare that:

TYPE OF DECLARATION

This declaration is for a national stage of PCT application.

INVENTORSHIP IDENTIFICATION

My residence, post office address and citizenship are as stated below, next to my name. I believe that I am an original, first and joint inventor of the subject matter that is claimed, and for which a patent is sought on the inventor entitled:

TITLE OF INVENTION

Improved Live Attenuated Viruses for Use as Vectors or Vaccines

SPECIFICATION IDENTIFICATION

The specification was filed on April 9, 1999, as International Application Number PCT/US99/07854.

ACKNOWLEDGMENT OF REVIEW OF PAPERS AND DUTY OF CANDOR

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

REQUEST FOR CHANGE IN AGENT'S ADDRESS

The correspondence address of Applicants' undersigned agent has changed.

Accordingly, pursuant to PCT Rule 92 bis, Applicants request that the International Bureau records the change of the agent's address to the following:

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Applicants further request the International Bureau to notify all Offices and PCT Authorities interested in the change accordingly.

In the event a fee is required and not enclosed, the Commissioner is authorized to charge any fee to the account of the undersigned attorneys. Account No. 50-1089.

Respectfully Submitted,

SAUL, EWING, REMICK & SAUL, L.L.P.

Janet E. Reed, Ph.D.

PTO Registration No. 36,252

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DECLARATION

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

SIGNATURE(S)

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